PCT

WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau



INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification 5: WO 91/09882 (11) International Publication Number: A1 C08F 4/76, 4/603, 10/00 11 July 1991 (11.07.91) (43) International Publication Date: (74) Agents: KURTZMAN, Myron, B. et al.; Exxon Chemical Company, 5200 Bayway Drive, Baytown, TX PCT/US90/07669 (21) International Application Number: (22) International Filing Date: 27 December 1990 (27:12.90)

US

(30) Priority data: 2 January 1990 (02.01.90) 459,921

(71) Applicant: EXXON CHEMICAL PATENTS INC. [US/ US]; 1900 East Linden Avenue, Linden, NJ 07036-0710

(72) Inventors: HLATKY, Gregory, George; 1114 Indian Autumn, Houston, TX 77062 (US). UPTON, David, Jonathan; 706 Glenmore, Pasadena, TX 77503 (US). TURN-ER, Howard, William; 303 Elder Glen, Webster, TX 77598 (US),

77522-5200 (US).

(81) Designated States: AT (European patent), AU, BE (Europesignated States: AI (European patent), AU, BE (European patent), BR, CA, CH (European patent), DE (European patent), DK (European patent), ES (European patent), FI, FR (European patent), GB (European patent), GR (European patent), HU, IT (European patent), JP, KR, LU (European patent), NL (European patent NO, SE (European patent), SU.

Published

With international search report. Before the expiration of the time limit for amending the claims and to be republished in the event of the receipt of amendments.

(54) Title: SUPPORTED IONIC METALLOCENE CATALYSTS FOR OLEFIN POLYMERIZATION

(57) Abstract

A supported catalyst for olefin polymerization comprising a Group IV-B metallocene component and an ionic activator component comprising a cation capable of donating a proton and a labile bulky anion having a plurality of lipophilic radicals so that the anion is sterically hindered from covalently bonding with a cation produced from the Group IV-B metal of the metallo-

FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

AT	Austria	ES	Spain	MG	Madagascar
AU.	Australia .	FI	Finland	ML	Mali
BB	Barbados	FR	France	MN	Mongolia
BE	Belgium	GA	Gabon	MR	Mauritania
BP	Burkina Faso	GB	United Kingdom	MW	Malawi
BG	Bulgaria	GN	Guinca	NL	Netherlands
BJ	Benin	GR	Greece	NO	Norway
BR	Brazil	HU	Hungary	PL	Poland
CA	Canada	IT	Italy	RO	Romania
CF	Central African Republic	JP	Japan	SD	Sudan
CG	Congo	KP	Democratic People's Republic	SE	Sweden
CH	Switzerland		of Korea	SN	Senegal
Ci	Côte d'Ivoire	KR	Republic of Korea	SU	Soviet Union
CM	Cameroon	Lì	Liechtenstein	TD	Chad
cs	Czechoslovakia	LK	Sri Lanka	TG	Togo
DE	Germany	LU	Luxembourg	us	United States of America
DY	Denmark	MC	Manne		

- 1 -

FOR OLEFIN POLYMERIZATION

Background of the Invention

1. Field of the Invention

1 The invention relates to supported catalysts for 2 polymerization of olefins including gas or slurry phase 3 polymerization of olefins, diolefins, cyclic olefins and 4 acetylenically unsaturated monomers. These catalysts, which may be 5 supported on known catalyst supports, include a Group IV-B metal 6 metallocene compound and an ionic activator compound. While the 7 homogeneous variant of this catalyst system has been previously 8 disclosed in copending U.S. Serial No. 133,480, a supported form of the catalyst system has not heretofore been produced. The supported 9 10 catalyst, suitable for use in gas or slurry phase olefin polymerization, provides a polymer product having a narrower particle size distribution and higher bulk density than achievable with the 13 homogeneous catalyst system. Furthermore, the use of the supported

catalyst in the gas phase results in greatly reduced reactor fouling as compared to the unsupported or homogeneous variant. By employing suitably sized supports, the catalyst system can be employed in solution and high pressure polymerization processes.

5

2. Background

Ziegler-Natta type catalysts for the polymerization of
olefins are well known. The traditional Ziegler-Natta type systems
comprise a metal halide activated to a catalyst species by reaction
with a metal alkyl cocatalyst, particularly an aluminum alkyl
cocatalyst. The activation of these traditional Ziegler-Natta
catalysts generates a variety of different active sites. As a
consequence of this non-uniformity of the active sites, the catalysts
produce polymer products of broad molecular weight distribution
(MWD). Furthermore, the copolymer products exhibit broad composition
distribution (CD), poor comonomer incorporation and blocky sequence
distribution.

Recently it has been found that active catalysts are formed when a bis(cyclopentadienyl) compound of the Group IV-B metals, in particular zirconium and hafnium, is activated by an alumoxane. The metallocene-alumoxane catalysts whether homogeneous or supported generally possess high activity and are more versatile than conventional Ziegler-Natta catalysts in that they may be effectively used to produce a variety of polymer products including, for example, high density linear polyethylene (HDPE), linear low density polyethylene (LLDPE), ethylene-propylene copolymer (EP), non-crystalline polypropylene and crystalline polypropylene. The metallocene-alumoxane catalysts also offer the significant advantage over the traditional Ziegler-Natta catalysts of being able to produce polymers with narrow MWD.

30 While the metallocene-alumoxane catalysts do offer 31 significant advantages over the traditional Ziegler-Natta catalysts, 32 they nevertheless have limitations in practical commercial 33 applications. These limitations include the relatively high cost of 34 the alumoxane cocatalysts. Alumoxane is also air sensitive and 2

4

difficult to manipulate. Furthermore, the metallocene-alumoxane catalysts, while producing a narrow MMD polymer product, have a limited capability to produce high molecular weight polymers or polymers having a high comonomer content.

5 Copending U.S. Patent Application Serial No. 133,480, also 6 published as European Patent Application 277,004 which is hereby incorporated by reference, describes a further advance in metallocene 7 catalysts: a new metallocene catalyst which does not require either 9 an alkyl aluminum or an alumoxane as an activator. The Group IV-B metallocene catalyst is prepared as a reaction product of a Group 10 IV-B metal metallocene compound and an ionic activator compound. The ionic activator comprises a cation having a donatable proton and a labile, bulky anion which is a single coordination complex having a plurality of lipophilic radicals covalently coordinated to and shielding a central charge-bearing metal or metalloid atom, the bulk of said anion being such that upon reaction of the activator cation donatable proton with a proton reactable substituent of a bis(cyclopentadienyl) Group IV-B metal compound to form a Group IV-B metal cation, the anion of the activator is sterically hindered from covalently coordinating to the Group IV-B metal cation. Hence, as described in the copending application, an active catalytic species 22 of a metallocene is formed, namely an ionic pair comprising a metallocene transition metal cation paired with a noncoordinating anion of the activator component.

The new metallocene catalyst system (hereafter referred to as an "ionic metallocene catalyst") eliminates the need for an expensive alumoxane activator. The ionic metallocene catalyst also offers other advantages over the metallocene-alumoxane catalysts such as permitting the production of polyolefin products of narrow MWD and of significantly higher weight average molecular weight at high rates of catalytic activity while also permitting better incorporation of comonomers and the control of the chain end chemistry of the polymer The new ionic metallocene catalyst of the copending application is, however, a homogeneous catalyst and generally can not be practically used for gas phase polymerization. The use of a

supported catalyst offers the possibility of gas phase 1 compatibility. Control of the particle size distribution of the 2 polymeric product in the various polymerization processes eliminates or reduces the extent of reactor fouling. 5 Supported catalysts for olefin polymerization are well known in the art. These catalysts offer, among others, the advantages of 7 being usable in gas or slurry phase reactors allowing the control of polymer particle size and thereby the control of product bulk density. Gas phase reactors also eliminate the need for a solvent and the equipment for solvent handling and separation. However, the known Ziegler-Natta olefin polymerization supported catalysts also present disadvantages which include broad MWD and composition distribution (CD), inefficient incorporation of comonomers, poor sequence distribution and, in the case of lower activity catalysts, 15 the need for a product deashing step. 16 Supported metallocene-alumoxane catalysts for olefin 17 polymerization are described in U.S. Patent 4,701,432 of Welborn. 18 These supported metallocene-alumoxane catalysts are obtained by 19 reacting a metallocene and an alumoxane in the presence of the solid 20 support material. The supported catalyst may then be employed either 21 as the sole catalyst component or may be employed in combination with 22 an organometallic cocatalyst. The supported metallocene-alumoxane 23 catalyst, however, still produces polymers of generally lower

It would be desirable to provide a supported catalyst for gas or slurry phase olefin polymerization that eliminates the need for either an alumoxane or an alkyl aluminum cocatalyst. It would be further desirable that such supported catalyst be capable of providing a polymer product having a high molecular weight, narrow MMD and CD, good comonomer incorporation, good sequence distribution, high bulk density and controlled particle size for ease of removal from the reactor.

24 molecular weight and comonomer incorporation than desired for certain

Summary

The invention provides a supported ionic metallocene 1 2 catalyst which is suitable for use in the polymerization of olefins including gas or slurry phase polymerization of olefins. The 3 heterogeneous catalyst, like its homogeneous counterpart disclosed in 4 our European Patent Application EP 277 004, permits the production of . 5 6 polyolefins of high molecular weight and narrow molecular weight 7 distribution (MWD) at high rates. Moreover, the polyolefin products 8 of the supported catalyst have a narrow composition distribution (CD) 9 and improved sequence distribution of comonomers as compared to the 10 products of prior art conventional supported Ziegler-Natta catalysts. 11 The possibility of producing a supported catalyst was 12 surprising since it would have been predicted that the reaction of ionic catalyst as described in the copending application with a Lewis base such as is present on a metal oxide surface would result in 15 catalyst deactivation. This invention is even more surprising since aluminum alkyls are not present during catalyst preparation and 17 furthermore the polymer products are similar to that obtained with 18 the unsupported catalyst. 19 The supported ionic metallocene catalyst of this invention 20 comprises the ionic metallocene catalyst and a suitable support 21 material. The metallocene component of the ionic metallocene 22 catalyst may be selected from a bis(cyclopentadienyl) derivative of a Group IV-B (Periodic Table of Elements, published and copyrighted by CRC Press, Inc., 1984) metal compound containing at least one ligand which will combine with an activator component or at least a portion 26 thereof such as a cation portion thereof. The activator component is an ionic compound comprising a cation which will irreversibly react 28 with at least one ligand contained in said Group IV-B metal compound 29 (metallocene component) and an anion which is a single coordination 30 complex comprising a plurality of lipophilic radicals covalently 31 coordinated to and shielding a central formally charge-bearing metal 32 or metalloid atom, which anion is bulky, labile and stable to any 33 reaction involving the cation of the activator component. The 34 charge-bearing metal or metalloid may be any metal or metalloid

```
1 capable of forming a coordination complex which is not hydrolyzed by
    aqueous solutions. Upon combination of the metallocene component and
2
    activator component, the cation of the activator component reacts
3
    with one of the ligands of the metallocene component, thereby
    generating an ion pair consisting of a Group IV-B metal cation with a
5
    formal coordination number of 3 and a valence of +4 and the
    aforementioned anion, which anion is compatible with and
7
    noncoordinating toward the metal cation formed from the metallocene
8
    component. The anion of the activator compound must be capable of
9
10 stabilizing the Group IV-B metal cation complex without interfering
11 with the ability of the Group IV-B metal cation or its decomposition
12 product to function as a catalyst and must be sufficiently labile to
13 permit displacement by an olefin, diolefin or an acetylenically
14 unsaturated monomer during polymerization.
             Either the ionic metallocene catalyst or both its components
15
16 will be contacted with an inorganic or organic solid support
17 material, either thermally or chemically dehydrated before such
18 contact, to form the supported ionic metallocene catalyst of this
19 invention.
             The method for preparing these supported ionic catalyst
20
21 comprises the steps of,
22
             (a) combining, in a solvent or diluent
23
                  (i) at least one metallocene component comprising a
24 bis(cyclopentadienyl) metal compound containing at least one ligand
25 capable of reacting with a proton, said metal being selected from
26 Group IV B metals,
27
                  (ii) at least one activator component comprising a
28 cation capable of donating a proton and an anion, said anion being a
29 single coordination complex comprising a plurality of lipophilic
30 radicals covalenty coordinated to and shielding a central
31 charge-bearing metal or metalloid atom, said anion being bulky,
32 labile and capable of stabilizing the metal cation formed as a result
33 of reaction between the two, and
34
                  (iii) a catalyst support material suspended in a
```

35 suitable solvent, and

3

4

5

6

(b) recovering a supported catalyst product as a free-flowing solid or slurry.

The supported catalyst of this invention will polymerize olefins, diolefins and/or acetylenically unsaturated monomers either alone or in combination with other olefins and/or other unsaturated monomers in liquid, slurry or gas phase reactions.

In general, catalysts can be selected so as to produce the polymer products which will be free of certain trace elements generally found in polymers produced with Ziegler-Natta type catalysts such as aluminum, magnesium, chloride and the like. The polymer products produced with the catalysts of this invention should, then, have a broader range of applications than polymers produced with either the more conventional Ziegler-Natfa type

14 catalysts containing a metal alkyl, such as an aluminum alkyl, or the 15 metallocene-alumoxane catalysts.

16 Detailed Description of the Preferred Embodiments

The present invention relates to supported catalysts
eminently suited for use in various polymerization processes
including gas or slurry phase polymerization of olefins. The
heterogeneous catalyst includes a metallocene of the Group IV-B
transition metals, an ionic activator compound and a catalyst
support. European Patent Application EP 277,004 incorporated by
reference, describes the homogeneous variant of the Group IV-B ionic
metallocene catalyst which is prepared as the reaction product of a
Group IV-B metal metallocene compound and an ionic activator compound.

The Metallocene Component

The Group IV-B metal compounds, particularly the titanium, zirconium and hafnium compounds, useful as first compounds in the ionic metallocene catalyst system employed in the process of this invention are the bis(cyclopentadienyl) derivatives of titanium, zirconium or hafnium. In general, such useful titanium, zirconium and hafnium compounds may be represented by the following general formulae:

1. (A-Cp)MX₁X₂

2 2. (A-Cp)MX'¸X'¸

3 3. (A-C_D)ML

4. (Cp*)(CpR)MX,

5 wherein: M is a metal selected from the Group consisting of titanium

6 (Ti), zirconium (Zr) and hafnium (Hf); (A-Cp) is either (Cp)(Cp*) or

7 Cp-A'-Cp* and Cp and Cp* are the same or different substituted or

g unsubstituted cyclopentadienyl radicals, and wherein A' is a covalent

9 bridging group containing a Group IV-A element; L is an olefin,

10 diolefin or aryne ligand; X_1 and X_2 are, independently,

11 selected from the group consisting of hydride radicals, hydrocarbyl

12 radicals having from 1 to about 20 carbon atoms,

13 substituted-hydrocarbyl radicals, wherein 1 or more of the hydrogen

14 atoms are replaced with a halogen atom, having from 1 to about 20

15 carbon atoms, organo-metalloid radicals comprising a Group IV-A

16 element wherein each of the hydrocarbyl substituents contained in the

17 organo-portion of said organo-metalloid, independently, contain from

18 1 to about 20 carbon atoms and the like; X'_1 and X'_2 are

19 joined and bound to the metal atom to form a metallacycle, in which

20 the metal, X'_1 and X'_2 form a hydrocarbocyclic ring

21 containing from about 3 to about 20 carbon atoms; and R is a

22 substituent, preferably a hydrocarbyl substituent, having from 1 to

23 20 carbon atoms, on one of the cyclopentadienyl radicals which is

24 also bound to the metal atom. Each carbon atom in the

25 cyclopentadienyl radical may be, independently, unsubstituted or

26 substituted with the same or a different radical selected from the

27 Group consisting of hydrocarbyl radicals, substituted-hydrocarbyl

28 radicals wherein one or more hydrogen atoms is replaced by a halogen

29 atom, hydrocarbyl-substituted metalloid radicals wherein the

30 metalloid is selected from Group IV-A of the Periodic Table of the

31 Elements, halogen radicals and the like. Suitable hydrocarbyl and

- 1 substituted-hydrocarbyl radicals which may be substituted for at
- 2 least one hydrogen atom in the cyclopentadienyl radical will contain
- 3 from 1 to about 20 carbon atoms and include straight and branched
- 4 alkyl radicals, cyclic hydrocarbon radicals, alkyl-substituted cyclic
- 5 hydrocarbon radicals, aromatic radicals and alkyl-substituted
- 6 aromatic radicals. Similarly, and when X, and/or X, is a
- 7 hydrocarbyl or substituted-hydrocarbyl radical, each may,
- s independently, contain from 1 to about 20 carbon atoms and be a
- 9 straight or branched alkyl radical, a cyclic hydrocarbyl radical, an
- 10 alkyl-substituted cyclic hydrocarbyl radical, an aromatic radical or
- 11 an alkyl-substituted aromatic radical. Suitable organometalloid
- 12 radicals include mono-, di- and trisubstituted organometalloid
- 13 radicals of Group IV-A elements wherein each of the hydrocarbyl
- 14 Groups contain from 1 to about 20 carbon atoms. Suitable
- 15 organometalloid radicals include trimethylsilyl, triethylsilyl,
- to ethyldimethylsilyl, methyldiethylsilyl, triphenylgermyl,
- 17 trimethylgermyl and the like.
- 18 Illustrative, but not limiting examples of
- 19 bis(cyclopentadienyl)zirconium compounds which may be used in the
- 20 preparation of the improved
- 21 (ethylcyclopentadienyl)(cyclopentadienyl) and
- 22 bis(ethylcyclopentadienyl)zirconium dihydride.
- 23 (propylcyclopentadienyl)(cyclopentadienyl) and
- 24 bis(propylcyclopentadienyl)zirconium dihydride,
- 25 (n-butylcyclopentadienyl)(cyclopentadienyl) and
- 26 bis(n-butylcyclopentadienyl)zirconium dihydride,
- 27 (t-butylcyclopentadienyl)(cyclopentadienyl) and
- 28 bis(t-butylcyclopentadienyl)zirconium dihydride,
- 29 (cyclohexylmethylcyclopentadienyl)(cyclopentadienyl) and
- 30 bis(cyclohexylmethylcyclopentadienyl)zirconium dihydride.
- 31 (benzylcyclopentadienyl)(cyclopentadienyl) and
- 32 bis(benzylcyclopentadienyl)zirconium dihydride,
- 33 (diphenylmethylcyclopentadienyl)(cyclopentadienyl) and
- 34 bis(diphenylmethylcyclopentadienyl)zirconium dihydride and the like;
- 35 (polyhydrocarbyl-substituted-cyclopentadienyl) zirconium compounds

- 1 such as (dimethylcyclopentadienyl) (cyclopentadienyl) and
- bis(dimethylcyclopentadienyl) zirconium dimethyl,
- 3 (trimethylcyclopentadienyl) (cyclopentadienyl) and
- 4 bis(trimethylcyclopentadienyl) zirconium dimethyl,
- 5 (tetramethylcyclopentadienyl) (cyclopentadienyl) and
- 6 bis(tetramethylcyclopentadienyl) zirconium dimethyl.
- 7 (permethylcyclopentadienyl) (cyclopentadienyl) and
- 8 bis(permethylcyclopentadienyl) zirconium dimethyl.
- 9 (ethyltetramethylcyclopentadienyl) (cyclopentadienyl) and
- 10 bis(ethyltetramethylcyclopentadienyl) zirconium dimethyl.
- 11 (indenyl)(cyclopentadienyl) and bis(indenyl)zirconium dimethyl,
- 12 (dimethylcyclopentadienyl) (cyclopentadienyl) and
- 13 bis(dimethylcyclopentadienyl) zirconium dihydride.
- 14 (trimethylcyclopentadienyl) (cyclopentadienyl) and
- 15 bis(trimethylcyclopentadienyl) zirconium dihydride.
- 16 (tetramethylcyclopentadienyl) (cyclopentadienyl) and
- 17 bis(tetramethylcyclopentadienyl)zirconium dihydride.
- 18 (permethylcyclopentadienyl) (cyclopentadienyl) and
- 19 bis(permethylcyclopentadienyl)zirconium dihydride.
- 20 (ethyltetramethylcyclopentadienyl) (cyclopentadienyl) and
- 21 bis(ethy)tetramethy)cyclopentadieny))zirconium dihydride.
- 22 (indenyl)(cyclopentadienyl) and bis(indenyl)zirconium dihydride
- 23 (propylcyclopentadienyl) (cyclopentadienyl) and
- 24 bis(propylcyclopentadienyl)zirconium dihydride,
- 25 (n-butylcyclopentadienyl) (cyclopentadienyl) and
- 26 bis(n-butylcyclopentadienyl)zirconium dihydride.
- 27 (t-butylcyclopentadienyl) (cyclopentadienyl) and
- 28 bis(t-butylcyclopentadienyl)zirconium dihydride.
- 29 (cyclohexylmethylcyclopentadienyl) (cyclopentadienyl) and
- 30 bis(cyclohexylmethylcyclopentadienyl)zirconium dihydride.
- 31 (benzylcyclopentadienyl) (cyclopentadienyl) and
- 32 bis(benzylcyclopentadienyl)zirconium dihydride.
- 33 (diphenylmethylcyclopentadienyl) (cyclopentadienyl) and
- 34 bis(diphenylmethylcyclopentadienyl)zirconium dihydride and the like:
- 35 (metal hydrocarbyl-substituted cyclopentadienyl)zirconium compounds

- i such as (trimethylsilylcyclopentadienyl)(cyclopentadienyl) and
- 2 bis(trimethylsilylcyclopentadienyl)zirconium dimethyl,
- 3 (trimethylgermylcyclopentadienyl)(cyclopentadienyl) and
- 4 bis(trimethylgermylcyclopentadienyl)zirconium dimethyl,
- 5 (trimethylstannylcyclopentadienyl)(cyclopentadienyl) and
- 6 bis(trimethylstannylcyclopentadienyl)zirconium dimethyl,
- 7 (trimethylplumbylcyclopentadienyl)(cyclopentadienyl) and
- 8 bis(trimethylplumbylcyclopentadienyl)zirconium dimethyl,
- 9 (trimethylsilylcyclopentadienyl)(cyclopentadienyl) and
- 10 bis(trimethylsilylcyclopentadienyl)zirconium dihydride,
- 11 (trimethylgermylcyclopentadienyl)(cyclopentadienyl) and
- 12 bis(trimethylgermylcyclopentadienyl)zirconium dihydride,
- 13 (trimethylstannylcyclopentadienyl)(cyclopentadienyl) and
- 14 bis(trimethylstannylcyclopentadienyl)zirconium dihydride,
- 15 (trimethylplumbylcyclopentadienyl)(cyclopentadienyl) and
- 16 bis(trimethylplumblycyclopentadienyl)zirconium dihydride and the
- 17 like; (halogen-substituted-cyclopentadienyl) zirconium compounds such
- 18 as (trifluoromethylcyclopentadienyl)(cyclopentadienyl) and
- 19 bis(trifluoromethylcyclopentadienyl)zirconium dimethyl
- 20 (trifluoromethylcyclopentadienyl)(cyclopentadienyl) and
- 21 bis(trifluoromethylcyclopentadienyl)zirconium dihydride and the like;
- 22 silyl-substituted bis(cyclopentadienyl) zirconium compounds such as
- 23 bis(cyclopentadienyl) (trimethylsilyl)(methyl)zirconium,
- 24 bis(cyclopentadienyl) (triphenylsilyl)(methyl)zirconium,
- 25 bis(cyclopentadienyl) [tris(dimethylsilyl)silyl](methyl)zirconium,
- 26 bis(cyclopentadienyl)[bis(mesityl)silyl](methyl)zirconium,
- 27 bis(cyclopentadienyl)(trimethylsilyl)trimethylsilylmethyl) zirconium,
- 28 bis(cyclopentadienyl) (trimethylsilylbenzyl) and the like:
- 29 (bridged-cyclopentadienyl)zirconium compounds such as methylene
- 30 bis(cyclopentadienyl)zirconium dimethyl, ethylene
- 31 bis(cyclopentadienyl)zirconium dimethyl, dimethylsilyl
- 32 bis(cyclopentadienyl)zirconium dimethyl, methylene
- 33 bis(cyclopentadienyl)zirconium dihydride, ethylene
- 34 bis(cyclopentadienyl)zirconium dihydride and dimethylsilyl
- 35 bis(cyclopentadienyl)zirconium dihydride and the like; zirconacycles

- such as bis(pentamethylcyclopentadienyl) zirconacyclobutane,
- bis(pentamethylcyclopentadienyl) zirconacyclopentane.
- 3 bis(cyclopentadienyl)zirconaindane.
- 4 1-bis(cyclopentadienyl)zircona-3-dimethylsila-cyclobutane and the
- 5 like; olefin, diolefin and aryne ligand substituted
- 6 bis(cyclopentadienyl)zirconium compounds such as
- 5 bis(cyclopentadienyl) (1,3-butadiene)zirconium, bis(cyclopentadienyl)
- 8 (2,3-dimethyl-1,3-butadiene)zirconium,
- 9 bis(pentamethylcyclopentadienyl)(benzyne)zirconium and the like;
- 10 (hydrocarbyl)(hydride) bis(cyclopentadienyl)zirconium compounds such
- 11 as bis(pentamethy)cyclopentadieny))zirconium (pheny))(hydride),
- 12 bis(pentamethylcyclopentadienyl)zirconium (methyl)(hydride) and the
- 13 like; and bis(cyclopentadienyl) zirconium compounds in which a
- 14 substituent on the cyclopentadienyl radical is bound to the metal
- 15 such as (pentamethylcyclopentadienyl)
- 16 (tetramethylcyclopentadienylmethylene) zirconium hydride.
- 17 (pentamethylcyclopentadienyl)
- 18 (tetramethylcyclopentadienylmethylene)zirconium phenyl and the like.
- A similar list of illustrative bis(cyclopentadienyl) hafnium
- 20 and bis(cyclopentadienyl)titanium compounds could be made, but since
- 21 the lists would be nearly identical to that already presented with
- 22 respect to bis(cyclopentadienyl)zirconium compounds, such lists are
- 23 not deemed essential to a complete disclosure. Other
- 24 bis(cyclopentadienyl)hafnium compounds and other
- 25 bis(cyclopentadienyl)titanium compounds as well as other
- 26 bis(cyclopentadienyl)zirconium compounds which are useful in the
- 27 catalyst compositions of this invention will, of course, be apparent
- 28 to those skilled in the art.

The Activator Component

- 29 Compounds useful as an activator component in the
- 30 preparation of the catalyst of this invention will comprise a cation,
- 31 which is a Bronsted acid capable of donating a proton, and a
- 32 compatible noncoordinating anion containing a single coordination
- 33 complex comprising a charge-bearing metal or metalloid core, which

anion is relatively large (bulky), capable of stabilizing the active catalyst species (the Group IV-B cation) which is formed when the two compounds are combined and said anion will be sufficiently labile to be displaced by olefinic, diolefinic and acetylenically unsaturated substrates or other neutral Lewis bases such as ethers, nitriles and Ś the like. As indicated above, any metal or metalloid capable of forming a coordination complex which is stable in water may be used 7 or contained in the anion of the second compound. Suitable metals, then, include, but are not limited to, aluminum, gold, platinum and the like. Suitable metalloids include, but are not limited to, boron, phosphorus, silicon and the like. Compounds containing anions 12 which comprise coordination complexes containing a single metal or 13 metalloid atom are, of course, well known and many, particularly such 14 compounds containing a single boron atom in the anion portion, are 15 available commercially. In light of this, salts containing anions 16 comprising a coordination complex containing a single boron atom are 17 preferred. In general, the activator compounds useful in the 18

19 preparation of the catalysts of this invention may be represented by

5. $[(L'-H)^+]^{d+}[(H')^{m+}Q_1Q_2...Q_n]^{d-}$

Wherein:

22

20 the following general formula:

L' is a neutral Lewis base; H is a hydrogen atom; [L'-H] is a Bronsted acid; M' is a metal or metalloid

25 selected from the Groups subtended by Groups V-B to V-A of the

26 Periodic Table of the Elements; i.e., Groups V-B, VI-B, VII-B,

27 VIII-B, I-B, II-B, III-A, IV-A, and V-A;

 Q_1 to Q_n are selected, independently, from the Group consisting of hydride radicals, dialkylamido radicals, alkoxide and anyloxide radicals, hydrocarbyl and substituted-hydro-carbyl radicals and organometalloid radicals and any one, but not more than one, of Q_1 to Q_n may be a halide radical — the remaining Q_1 to Q_n being, independently, selected from the foregoing radicals:

m is an integer from 1 to 7; n is an integer from 2 to 8; 1 2 and n - m = d. The preferred activator compounds comprising boron may be represented by the following general formula: 6. [L'H] [BAr, Ar, X, X,] 6 Wherein: L' is a neutral Lewis base; H is a hydrogen atom: [L'-H] ts a Bronsted acid; B is boron in a valence state 10 of 3; Ar_1 and Ar_2 are the same or different aromatic or 11 substituted-aromatic hydrocarbon radicals containing from about 6 to 12 about 20 carbon atoms and may be linked to each other through a 13 stable bridging group; and X_2 and X_4 are radicals selected, 14 independently, from the group consisting of hydride radicals, halide 15 radicals, with the proviso that only X_2 or X_4 will be halide 16 at the same time, hydrocarbyl radicals containing from 1 to about 20 17 carbon atoms, substituted-hydrocarbyl radicals, wherein one or more 18 of the hydrogen atoms is replaced by a halogen atom, containing from 19 1 to about 20 carbon atoms, hydrocarbyl-substituted metal 20 (organometalloid) radicals wherein each hydrocarbyl substitution 21 contains from 1 to about 20 carbon atoms and said metal is selected 22 from Group IV-A of the Periodic Table of the Elements and the like. 23 In general, Ar, and Ar, may, independently, be any 24 aromatic or substituted-aromatic hydrocarbon radical containing from 25 about 6 to about 20 carbon atoms. Suitable aromatic radicals 26 include, but are not limited to, phenyl, naphthyl and anthracenyl 27 radicals. Suitable substituents include, but are not necessarily 28 limited to, hydrocarbyl radicals, organometalloid radicals, alkoxy

radicals, alkylamido radicals, fluoro and fluorohydrocarbyl radicals and the like such as those useful as X_3 and X_4 . The substituent may be ortho, meta or para, relative to the carbon atoms

32 bonded to the boron atom. When either or both X_2 and X_4 are

- 1 a hydrocarbyl radical, each may be the same or a different aromatic
- 2 or substituted-aromatic radical as are Ar_1 and Ar_2 , or the
- 3 same may be a straight or branched alkyl, alkenyl or alkynyl radical
- 4 having from 1 to about 20 carbon atoms, a cyclic hydrocarbon radical
- 5 having from about 5 to about 8 carbon atoms or an alkyl-substituted
- 6 cyclic hydrocarbon radical having from about 6 to about 20 carbon
- 7 atoms. X_2 and X_4 may also, independently, be alkoxy or
- 8 dialkylamido radicals wherein the alkyl portion of said alkoxy and
- 9 dialkylamido radicals contain from 1 to about 20 carbon atoms,
- 10 hydrocarbyl radicals and organometalloid radicals having from 1 to
- 11 about 20 carbon atoms and the like. As indicated above, Ar, and
- 12 Ar may be linked to each otherN Similarly, either or both of
- 13 Ar and Ar could be linked to either X_3 or X_4 .
- 14 Finally, X_a or X_4 may also be linked to each other through a
- 15 suitable bridging group.
- 16 Illustrative, but not limiting, examples of boron compounds
- 17 which may be used as an activator component in the preparation of the
- 18 improved catalysts of this invention are trialkyl-substituted
- 19 ammonium salts such as triethylammonium tetra(phenyl)boron.
- 20 tripropylammonium tetra(phenyl)boron, tri(n-butyl)ammonium
- 21 tetra(phenyl)boron, trimethylammonium tetra(p-tolyl)boron,
- 22 trimethylammonium tetra(o-tolyl)boron, tributylammonium
- 23 tetra(pentafluorophenyl)boron, tripropylammonium
- 24 tetra(o,p-dimethylphenyl)boron, tributylammonium
- 25 tetra(m,m-dimethylphenyl)boron, tributylammonium
- 26 tetra(p-tri-fluoromethylphenyl)boron, tributylammonium
- 27 tetra(pentafluorophenyl)boron, tri(n-butyl)ammonium
- 28 tetra(o-tolyl)boron and the like; N.N-dialkyl anilinium salts such as
- 29 N.N-dimethylanilinium tetra(phenyl)boron, N.N-diethylanilinium
- 30 tetra(phenyl)boron, N,N-2,4,6-pentamethylanilinium tetra(phenyl)boron
- 31 and the like; dialkyl ammonium salts such as di(i-propyl)ammonium
- 32 tetra(pentafluorophenyl)boron, dicyclohexylammonium
- 33 tetra(phenyl)boron and the like; and triaryl phosphonium salts such
- 34 as triphenylphosphonium tetra(phenyl)boron,
- 35 tri(methylphenyl)phosphonium tetra(phenyl)boron,

- tri(dimethylphenyl)phosphonium tetra(phenyl)boron and the like.
- 2 Suitable compounds containing other metals and metalloids
- which are useful as activator components can be usefully employed. 3
- In this regard, it should be noted that the foregoing list is not
- 5 intended to be exhaustive and that other useful boron compounds as
- well as useful compounds containing other metals or metalloids would
- be readily apparent to those skilled in the art from the foregoing 7
- general equations.

The Catalyst Support

- Typically, the support can be any of the known solid
- 10 catalyst supports, particularly porous supports, such as talc,
- 11 inorganic oxides, and resinous support materials such as
- 12 polyolefins. Preferably, the support material is an inorganic oxide
- 13 in particulate form.
- Suitable inorganic oxide materials which are desirably 14
- 15 employed in accordance with this invention include Group II-A, III-A,
- 16 IV-A or IV-B metal oxides. The most preferred catalyst support
- 17 materials include silica, alumina, and silica-alumina and mixtures
- 18 thereof. Other inorganic oxides that may be employed either alone or
- 19 in combination with the silica, alumina or silica-alumina are
- 20 magnesia, titania, zirconia, and the like. Other suitable support
- 21 materials, however, can be employed, for example, finely divided
- 22 polyolefins such as finely divided polyethylene.
- The metal oxides generally contain surface hydroxyl groups 23
- 24 which may react with and deactivate the ionic metallocene catalyst
- 25 when the catalyst is added to the slurried metal oxide support.
- 26 Therefore, it is preferred that the inorganic oxide support be
- 27 dehydrated prior to use, i.e. subjected to a thermal treatment in
- 28 order to remove water and reduce the concentration of the surface
- 29 hydroxyl groups. The treatment may be carried out in vacuum or while
- 30 purging with a dry inert gas such as nitrogen at a temperature of
- 31 about 100°C to about 1000°C, and preferably, from about 300°C to
- 32 about 800°C. Pressure considerations are not critical. The duration
- 33 of the thermal treatment can be from about 1 to about 24 hours.

1 However, shorter or longer times can be employed.

As an alternative method of dehydration of the metal oxide 2 support material, chemical dehydration can be advantageously 3 employed. Chemical dehydration converts all water and hydroxyl groups on the oxide surface to inert species. Useful chemical agents 5 are for example, chlorosilanes, such as trimethylchlorosilane, and 6 the like and alkyl aluminum reagents such as triethyl aluminum and 7 the like. The chemical dehydration is accomplished by slurrying the 8 inorganic particulate material, such as, for example, silica in an inert low boiling hydrocarbon, such as, for example, hexane. During the chemical dehydration reaction, the silica should be maintained in a moisture and oxygen-free atmosphere. To the silica slurry is then added a low boiling inert hydrocarbon solution of the chemical dehydrating agent. 14

The inorganic oxide support used in the preparation of the catalyst may be any particulate oxide or mixed oxide as previously described which has been thermally or chemically dehydrated such that it is substantially free of adsorbed moisture.

The specific particle size, surface area and pore volume of the inorganic oxide determine the amount of inorganic oxide that is desirable to employ in preparing the catalyst compositions, as well as affecting the properties of polymers formed with the aid of the catalyst compositions. These properties must frequently be taken into consideration in choosing an inorganic oxide for use in a particular aspect of the invention. In general, optimum results are usually obtained by the use of inorganic oxides having an average particle size in the range of about 0.1 to 600 microns, preferably about 0.3 to 80 microns; a surface area of about 50 to 1,000 square meters per gram, preferably about 100 to 400 square meters per gram; and a pore volume of about 0.5 to 3.5 cc per gram; preferably about 31 0.5 to 2 cc per gram.

Catalyst Preparation and Use

The supported ionic metallocene catalyst of this invention may be prepared by combining the metallocene component, the activator

- 1 component and the support in suitable solvents in one or more
- 2 steps.

A. Choice of Metallocene-Activator Pairs

In general, while most metallocene components identified above may be combined with most activator components identified above to produce an active olefin polymerization catalyst, it is desirable 5 for continuity of the polymerization operations that either the metal cation initially formed from the metallocene component or a decomposition product thereof be a relatively stable catalyst. It is also desirable that the anion of the activator compound be stable to 10 hydrolysis when an ammonium salt is used. Further, it is desirable 11 that the acidity of the activator component be sufficient, relative 12 to the metallocene component to facilitate the needed proton 13 transfer. In general, bis(cyclopentadienyl) metal compounds which 14 can be hydrolyzed by aqueous solutions can be considered suitable as metallocene components to form the catalysts described herein. The chemical reactions which occur in forming the catalyst 16 17 of this invention may, when a preferred, boron-containing compound is 18 used as the second component, be represented by a reference to the 19 general formulae set forth herein as follows:

- 20 7. $(A-Cp)HX_1X_2 + [L'-H]^+[BAr_1Ar_2X_3X_4]^- \rightarrow [(A-Cp)HX_1]^+[BAr_1Ar_2X_3X_4]^- + HX_2 + L' \text{ or } [(A-Cp)HX_2]^+[BAr_1Ar_2X_3X_4]^- + HX_1 + L'$
- 23 8. (A-Cp)HX'₁X'₂ + [L'-H]⁺[BAr₁Ar₂X₃X₄]⁻ 24 [(A-Cp)H(X'₁X'₂H)]⁺[BAr₁Ar₂X₃X₄]⁻ + L' or 25 [(A-Cp)H(X'₂X'₁H)]⁺[BAr₁Ar₂X₃X₄]⁻ + L'
- 26 9. (A-Cp)ML + [L'-H]⁺[BAr₁Ar₂X₃X₄]⁻ → [(A-Cp)M(LH)]⁺[BAr₁Ar₂X₃X₄]⁻ + L'
- 28 10. (Cp)(R-Cp*)MX₁] +[L'-H]⁺[BAr₁Ar₂X₃X₄] 29 [Cp(HR-Cp*)MX₁]⁺[BAr₁Ar₂X₃X₄] + L' or 30 [Cp(R-Cp*)M]⁺[BAr₁Ar₂X₃X₄] + HX₁ + L'

374

In the foregoing reaction equations the symbols have been previously defined. In general, the stability and rate of formation of the products in the foregoing reaction equations, particularly the 3 metal cation, will vary depending upon the choice of the solvent, the 4 acidity of the [L'-H]+ selected, the particular L', the anion, the 5 temperature at which the reaction is completed and the particular 6 bis(cyclopentadienyl) derivative of the metal selected. Generally, the initially formed ion-pair will be an active polymerization 8 catalyst and will polymerize α -olefins, diolefins, strained cyclic 9 10 olefins and acetylenically unsaturated monomers either alone or in combination with other monomers. In some cases, however, the initial metal cation will decompose to yield an active polymerization catalyst. With respect to the combination of the metallocene component 13 14 with the activator component to form a catalyst of this invention, it 15 should be noted that the two compounds combined for preparation of the 16 active catalyst must be selected to avoid transfer of a fragment of 17 the anion, particularly an aryl group, to the metal cation, thereby 18 forming a catalytically inactive species. This can be done by steric 19 hindrance, resulting from substitutions on the cyclopentadienyl carbon 20 atoms as well as substitutions on the aromatic carbon atoms of the 21 anion. It follows, then, that the metallocene components comprising 22 perhydrocarbyl-substituted cyclopentadienyl radicals could be 23 effectively used with a broader range of activator compounds than 24 could metallocene components comprising unsubstituted cyclopentadienyl 25 radicals. As the amount and size of the substitutions on the 26 cyclopentadienyl radicals are reduced however, more effective 27 catalysts are obtained with activator compounds containing anions 28 which are more resistant to degradation, such as those with 29 substituents on the ortho positions of the phenyl rings. Another 30 means of rendering the anion more resistant to degradation is afforded 31 by fluorine substitution, especially perfluoro-substitution, in the 32 anion. Fluoro-substituted stabilizing anions may, then, be used with 33 a broader range of metal compound (first components). Activators in 34 which the anions comprise pentafluorophenyl groups are preferred for 35 preparing ion-pair metallocene catalysts of this invention.

- It is preferred that the mole ratio of metallocene component
- to activator component be 1:1 or greater. In a reaction corresponding
- to general formula 1, when the anion comprises pentafluorophenyl
- groups, two structurally distinct forms of thermally stable ionic
- catalysts have been identified by NMR spectroscopy and are shown in
- general formulae 5 and 6, using tetrakis(pentafluorophenyl)boron as a
- representative anion:
- 11. $[(A-Cp)MX_1(L')][B(C_F_B)_4]$ 12. $[\{(A-Cp)MX_1\}_2(\mu-X_1)][B(C_F_B)_4]$
- 10 In the foregoing formulae, the symbols A-Cp, M, L', and X_1
- 11 correspond to the definitions set forth in general formulae 1-4. The
- 12 symbol "Y" indicates the ligand X1 bridges the two metal
- 13 centers. In both general formulae 5 and 6, NMR experiments indicate
- 14 that the fluorinated boron anion is completely non-coordinating.
- 15 When the molar ratio of metallocene component to activator component
- 16 is 1:1, L' weakly coordinates to an stabilizes the metallocene cation
- 17 when L' is an aniline derivative, for example N.N-dimethylaniline, to
- 18 give an ionic catalyst of general formula 5. When the molar ratio of
- 19 metallocene component to activator component is greater than 1:1 an X
- 20 group, for examples, a methyl group of an excess
- 21 $(A-Cp)M(CH_3)_2$ molecule when $(A-Cp)M(CH_3)_2$ is the
- 22 metallocene used, weakly coordinates to and stabilizes the
- 23 metallocene cation to give an ionic catalyst of general formula 6.

Catalyst Preparation

- The supported catalyst of this invention can be prepared by 24
- 25 combining the metallocene, activator components and support in one or
- 26 more suitable solvents or diluent. Suitable solvents and/or diluents
- 27 include, but are not necessarily limited to, straight and
- 28 branched-chain hydrocarbons such as isobutane, butane, pentane,
- 29 hexane, heptane, octane and the like; cyclic and
- 30 alicyclic hydrocarbons such as cyclohexane, cycloheptane,
- 31 methylcyclohexane, methylcycloheptane and the like; and aromatic and

3

5

19

alkyl-substituted aromatic compounds such as benzene, toluene, xylene and the like.

It is preferred that the catalyst components be handled in an inert, moisture-free, oxygen-free environment such as argon, nitrogen or helium because of the sensitivity of the catalyst components to moisture and oxygen.

In a preferred method, the metallocene and activator

components are combined in a first step in an aromatic solvent to

produce a solution the reaction product. This reaction may be carried

out in the temperature range -100° to about 300°C, preferably about 0°

to about 100°C. Holding times to allow for the completion of the

reaction may range from about 10 seconds to about 60 minutes depending

upon variables such as reaction temperature and choice of reactants.

The solution produced by combining the metallocene and activator components is then contacted with the support. The method of contact may vary, but it is preferred that the solution be added to a rapidly stirred slurry of the catalyst support in a hydrocarbon solvent, preferably an aliphatic solvent and especially pentane.

In another preferred method, in a first step, the activator

component is dissolved in an aromatic solvent with the support to produce a supported activator component. This reaction is carried out at a temperature sufficient to produce a homogeneous solution of the activator component, preferably between about 25°C to about 200°C.

The aromatic solvent is then removed to leave a free-flowing supported material. The supported material is then contacted with the metallocene component, preferably in an aliphatic solvent to produce the supported catalyst.

Regardless of the method, the active supported catalyst can be recovered by evaporation of the solvent to obtain a free-flowing solid or alternatively, the active supported catalyst can be maintained in its slurry state for direct use.

Contact temperatures may range from about 0° to about 100°C depending upon the solvents used. Contact times may vary from about 10 seconds to about 60 minutes, longer contact times than 60 minutes not providing any significant additional benefits.

2

3

35

In the preparation of the supported catalyst, the reagents should be combined to provide a catalyst concentration (metallocene and activator) on the support of from about 0.01 wt % to about 20 wt %, preferably about 1 wt % to about 5 wt % based upon the weight of the support.

5 In a most preferred embodiment of the present invention, 6 bis(cyclopentadienyl)zirconium dimethyl or bis(cyclopentadienyl)hafnium dimethyl will be reacted with 8 N,N-dimethylanilinium tetra(pentafluorophenyl)boron and then contacted 9 10 with an alumina, silica or silica-alumina support to produce the most preferred catalyst of the present invention. The metallocene and 11 activator components will be combined at a temperature within the 12 range -100° to about 300°C, preferably from about 0°C to about 100°C, 13 and preferably in an aromatic hydrocarbon solvent, most preferably 14 toluene. A nominal holding time within the range from about 10 15 seconds to about 60 minutes will be sufficient before the mixture is 16 contacted with a slurry of the dried support material in an aliphatic 17 solvent for a contacting period of from about 10 seconds to about 60 18 19 minutes.

20 In another preferred method, in a first step, N,N-dimethylanilinium tetra(pentafluorophenyl)boron is dissolved in an 21 aromatic solvent maintained at a temperature sufficient to dissolve 22 the activator component. The support is slurried therein and the 23 activator and support are reacted for about 1 minute to about 1 hour to produce a supported activator component. The aromatic solvent is 25 removed to leave a free-flowing support material containing the activator component. This supported activator component is then 27 contacted with bis(cyclopentadienyl)zirconium dimethyl or 28 bis(cyclopentadienyl)hafnium dimethyl, preferably in an aliphatic 29 solvent to produce the supported catalyst. 30

With either method, the active supported catalyst can be recovered by evaporation of the solvent to obtain a free-flowing solid or, alternately, the active supported catalyst can be maintained in a slurry state for direct use.

C. Catalyst Use

The supported ionic metallocene catalyst may be used to

polymerize α -olefins and acetylenically unsaturated monomers having 1 from 2 to about 18 carbon atoms and/or diolefins having from 4 to 2 about 18 carbon atoms either alone or in combination. The catalyst 3 may also be used to polymerize α -olefins, diolefins, strained cyclic 4 olefins and/or acetylenically unsaturated monomers in combination with 5 other unsaturated monomers. While the catalyst is active for this broad range of olefinic monomer feedstock, α -olefin polymerization 7 is preferred especially the homopolymerization of ethylene or the 8 copolymerization of ethylene with olefins having from 3 to 10 carbon 9 atoms. 10 In a preferred embodiment of the present invention, the 11 metallocene component will be a bis(cyclopentadienyl)-Group IV-B metal compound containing two, independently, substituted or unsubstituted 13 cyclopentadienyl radicals and one or two lower alkyl substituents and/or one or two hydride substituents and the activator component will be a tri-substituted ammonium salt of a fluorinated tetraphenyl boron anion. Each of the tri-substitutions in the ammonium cation will be the same or a different lower alkyl or aryl radical. By lower alkyl is meant an alkyl radical containing from 1 to 4 carbon atoms. 19 N,N-dimethylanilinium tetra(pentafluorophenyl)boron is particularly 20 preferred. Alumina, silica or silica-alumina will be the preferred 21 support for producing the supported ionic metallocene catalyst. 22 Certain of the catalysts of this invention, particularly 23 those based on hafnocenes - using the catalyst produced from the 24 reaction of bis(cyclopentadienyl)hafnium dimethyl and the tri-substituted ammonium salt of tetra(pentafluorophenyl)boron as an example - when used as described herein for the polymerization and copolymerization of α -olefins, diolefins, and/or acetylenically unsaturated monomers, in the absence of a chain transfer agent, can

28 Copolymerization of α-olefins, diolefins, and/or acetylenically
29 Unsaturated monomers, in the absence of a chain transfer agent, can
30 lead to the production of extremely high molecular weight polymers and
31 copolymers having relatively narrow molecular weight distributions.
32 In this regard, it should be noted that homopolymers and copolymers
33 having molecular weights up to about 2 x 10⁶ or higher and

molecular weight distributions within the range of about 1.5 to about 35 3 or greater can be produced with the catalysts of this invention.

35

```
Combinations of two or more ionic metallocene catalyst with the
    support can be employed in order to obtain broader MWD such as up to
   about 15 or greater. Alternatively, two or more separately supported
    catalysts can be used to obtain broad MWD polymers and copolymers.
5
             Supported catalysts of this invention containing a
   metallocene component which is either a pure enantiomer or the racemic
   mixture of two enantiomers of a rigid, chiral metallocene can
   polymerize prochiral olefins (propylene and higher \alpha-olefins) to
   crystalline polymers including syndiotactic and isotactic polymers.
10 Bis(cyclopentadienyl)metal compounds in which each of the
11 cyclopentadienyl radicals is substituted and containing a covalent
12 bridging group between the two cyclopentadienyl radicals are
13 particularly useful for isotactic polymerizations of this type.
14 Prochiral metallocenes, for example those based on complexes of
15 propyl-2-cyclopentadienyl-2-(1-fluorenyl) hafnium, can be used to
16 polymerize propylene or higher \alpha-olefins to syndiotactic polymers.
             The supported catalysts may be most usefully employed in gas
17
18 or slurry phase processes, both of which are known to those of skill
19 in the art. Thus, polymerizations using the invention supported
20 catalysts may be conducted by either of these processes, generally at
21 a temperature in the range of about 0°-160°C or even higher, and under
22 atmospheric, subatmospheric, or superatmospheric pressure conditions.
            A slurry polymerization process can utilize sub- or
23 ...
24 super-atmospheric pressures and temperatures in the range of
25 -80-250°C. In a slurry polymerization, a suspension of solid.
26 particulate polymer is formed in a liquid polymerization medium to
27 which ethylene, \alpha-olefin, diolefin, cyclic olefin or acetylenically
28 unsaturated comonomer, hydrogen and catalyst are added. Alkanes and
29 cycloalkanes, such as butane, pentane, hexane, or cyclohexane, are
30 preferred with C_4 to C_{10} alkanes especially preferred.
31 Preferred solvents also include liquid olefins which may act as
32 monomers or comonomers including ethylene, propylene, butadiene,
33 cyclopentene, 1-hexene, 3-methyl-1-pentene, 4-methyl-1-pentene,
34 1.4-hexadiene, 1-octene, 1-decene and the like.
```

A gas-phase polymerization process utilizes superatmospheric

- 1 pressure and temperatures in the range of about 50° 120°C.
- 2 Gas-phase polymerization can be performed in a stirred or fluidized
- 3 bed of catalyst and product particles in a pressure vessel adapted to
- 4 permit the separation of product particles from unreacted gases.
- 5 Thermostated ethylene, comonomer, hydrogen and an inert diluent gas
- 6 such as nitrogen can be introduced or recirculated so as to maintain
- 7 the particles at a temperature of 50° 120°C. Polymer product can be
- 8 withdrawn continuously or semi-continuously at a rate such as to
- 9 maintain a constant product inventory in the reactor. After
- 10 polymerization and deactivation of the catalyst, the product polymer
- 11 can be recovered by any suitable means. In commercial practice, the
- 12 polymer product can be recovered directly from the gas phase reactor,
- 13 freed of residual monomer with a nitrogen purge, and used without
- 14 further deactivation or catalyst removal. The polymer obtained can be
- 15 extruded into water and cut into pellets or other suitable comminuted
- 16 shapes. Pigments, antioxidants and other additives, as is known in
- 17 the art, may be added to the polymer.
- 18 While it is a characteristic of the invention supported
- 19 catalyst that the produced polymers have a narrow molecular weight
- 20 distribution, broad molecular weight distribution polymers may be
- 21 produced by using two or more metallocenes or two or more activators.
- The advantages of the instant invention will be more readily
- 23 appreciated by reference to the following illustrative, non-limiting
- 24 examples.

- 25 A supported catalyst was prepared by reacting
- 26 bis(cyclopentadienyl)hafnium dimethyl (30 mg) with
- 27 N.N-dimethylanilinium tetrakis(pentafluorophenyl)boron (60 mg) in
- 28 toluene (8 ml). This solution was added slowly to a suspension of
- 29 basic alumina (Brockman Activity I, dried overnight at 100°C under
- 30 vacuum; 2.0 g) in pentane (25 ml). The mixture was allowed to stir
- 31 for about 3 minutes at room temperature. The supernatant solution was
- 32 then decanted from the solid material and the solids reslurried with
- 33 fresh pentane. The supernatant solution was again decanted from the

- solids and the solids suspended in fresh pentane (ca. 30 ml). This suspension was transferred, under nitrogen, by means of a double-ended
- 3 needle into a 1 liter stainless-steel autoclave containing 400 ml of
- 4 dry, deoxygenated hexane and which had been previously flushed with
- 5 nitrogen. The autoclave was then pressured to 90 psig with ethylene
- 6 and stirred at 40°C. After 30 minutes, the autoclave was cooled,
- 7 vented and the contents separated by filtration. The yield of linear
- 8 granular, free-flowing polyethylene was 11.2 g. The polymer had a
- 9 weight-average molecular weight of 594,000 and a molecular weight
- 10 distribution of 2.15.

The supported catalyst of Example 1 was used to polymerize ethylene in a process similar to that of Example 1 except that the autoclave was pressured to 300 psig of ethylene. The yield of linear granular, free-flowing polyethylene was 48.5 g with a bulk density of 0.17 g/cc versus 0.07 g/cc when an unsupported, homogeneous catalyst is used.

Example 3

A supported catalyst was prepared by reacting
bis(cyclopentadienyl)zirconium dimethyl (20 mg) with
N,N-dimethylanilinium tetrakis(pentafluorophenyl)boron (60 mg) in
toluene (20 ml). This solution was added slowly to a suspension of
basic alumina (2.0 g) in pentane (25 ml). The mixture was allowed to
stir for about 3 minutes at room temperature. The supernatant
solution was decanted from the solid material and the solids
reslurried with fresh pentane. The supernatant solution was again
decanted from the solids and the solids were suspended in fresh
pentane (ca. 30 ml). This suspension was transferred, under nitrogen,
by means of a double-ended needle into a l liter stainless-steel
autoclave, previously flushed with nitrogen, containing 400 ml of dry,
deoxygenated hexane. The autoclave was then pressured to 400 psig
with ethylene and stirred at 40°C. After 30 minutes, the autoclave

- 1 yield of linear granular, free-flowing polyethylene was 4.7 g with a
- 2 weight-average molecular weight of 515,000 and a MWD of 1.74.

- A supported catalyst was prepared as in Example 1 except that
- 2.0 g of Davison 948 silica (dried at 800°C with a nitrogen purge) was
- 6 used in place of alumina. The catalyst was used to polymerize
- 7 ethylene in a process similar to that of Example 2. The yield of
- 8 linear granular, free-flowing polyethylene was 11.1 g with a
- 9 weight-average molecular weight of 1,384,000 and a MWD of 1.67.

Example 5

- A supported catalyst was prepared as in Example 4 except that
- 11 20 mg of bis(cyclopentadienyl)zirconium dimethyl was used in place of
- 12 bis(cyclopentadienyl)hafnium dimethyl. This catalyst was used to
- 13 polymerize ethylene in a process similar to that of Example 2. The
- 14 yield of linear granular, free-flowing polyethylene formed was 2.7 g
- 15 with a weight-average molecular weight of 759,000 and a MWD of 1.67.

Example 6

- A supported catalyst was prepared by reacting.
- 17 bis(cyclopentadienyl)hafnium dimethyl (45 mg) with
- 18 N,N-dimethylanilinium tetrakis(pentafluorophenyl)boron (90 mg) in
- 19 toluene (20 ml). This solution was added slowly to a suspension of
- 20 basic alumina (3.0 g) in pentane (25 ml). The mixture was allowed to
- 21 stir for about 3 minutes at room temperature. The solution was
- 22 decanted from the solid material and the solids reslurried with fresh
- 23 pentane. The supernatant solution was again decanted from the solids
- 24 and the solids were dried in an atmosphere of nitrogen. This
- 25 suspension was injected into a 1 liter stainless-steel autoclave,
- 26 previously flushed with nitrogen, containing 300 ml of propylene, by
- 27 washing 50 ml of propylene through a stainless-steel catalyst addition
- 28 tube containing the solid catalyst. The autoclave was then pressured
- 29 to 100 psig with ethylene and stirred at 50°C. After 15 minutes, the
- 30 autoclave was cooled and vented and the polymer product was isolated.

- 1 The yield of granular ethylene-propylene copolymer was 38 g. This
- 2 polymer, which contained 21 wt. % ethylene, had a weight-average
- 3 molecular weight of 423,000 and a molecular weight distribution of
- 4 2.88.

- 5 Ethylene and propylene was copolymerized in a manner similar
- 6 to Example 6, with the exceptions that the ethylene pressure was
- 7 increased to 200 psig and the autoclave was heated to 45°C. The yield
- 8 of granular ethylene-propylene copolymer was 57.1 g. This polymer.
- 9 which contained 52 wt. % ethylene, had a weight-average molecular
- 10 weight of 1,006,000 and a molecular weight distribution of 2.01.

Example 8

- A supported catalyst of this invention was prepared by
- 12 reacting rac-dimethylsilylbis(indenyl)hafnium dimethyl (29 mg) with
- 13 N,N-dimethylanilinium tetrakis(pentafluorophenyl)boron (45 mg) in
- 14 toluene (10 ml). This solution was added slowly to a suspension of
- 15 basic alumina (1.5 g) in pentane (25 ml). The mixture was allowed to
- 16 stir for about 3 minutes at room temperature. The supernatant
- 17 solution was decanted from the solid material and the solids
- 18 reslurried with fresh pentane. The supernatant solution was again
- 19 decanted from the solids and the solids were suspended in fresh
- 20 pentane (ca. 30 ml). This suspension was transferred under nitrogen,
- 21 by means of double-ended needle into a 1 liter stainless-steel
- 22 autoclave, previously flushed with nitrogen, containing 400 ml of dry.
- 23 deoxygenated hexane. Propylene (200 ml) was added to the autoclave.
- 24 which was stirred at 50°C. After 1.75 hours, the autoclave was cooled
- 25 and vented and the contents separated by filtration. The yield of
- 26 isotactic polypropylene was 9.5 g. This polymer had a melting point
- 27 of 127°C, as determined by differential scanning calorimetry, a weight
- 28 average molecular weight of 181,000 and a molecular weight
- 29 distribution of 1.95. Analysis by C-13 NMR spectroscopy indicated
- 30 that the polypropylene was 92% isotactic.

A l liter stainless-steel autoclave was charged with virgin polypropylene (14 mesh, 190 g) and heated to 85°C under a dynamic nitrogen purge. After cooling to 50°C, the catalyst of Example 1 (2.0 g) was added to the reactor from a stainless-steel catalyst addition tube using high-pressure nitrogen. After venting off the excess nitrogen pressure, ethylene was admitted to the stirred polymer bed to maintain a constant flow. The pressure inside the autoclave ranged from 10 to 35 psig. The maximum temperature in the autoclave reached 70°C. After 1 hour, the autoclave was cooled and vented and the polymer mixture isolated. The net increase in the polymer weight was 13 g. A portion of the polyethylene was sieved from the polypropylene stirring aid and was found to have a weight-average molecular weight 3 of 175,000 and a MMD of 2.53.

Example 10

Alumina (1.0 g), slurried in 10 ml of pentane, was treated for 15 minutes with 1 ml of a 1 M solution of triethylaluminum in the hexane. The support was separated from the liquid by decantation, washed twice with 10 ml of pentane and dried in vacuo.

N,N-dimethylanilinium tetrakis (pentafluorophenyl)boron (30 mg) was heated in toluene (25 ml) to produce a homogeneous solution. The warm solution was added dropwise to the support prepared as described above. The solvent was removed in vacuo to leave a free-flowing supported material.

This support material (1.0 g) was slurried in 10 ml of pentane. To this was added dropwise 5 ml of a pentane solution.

This support material (1.0 g) was slurried in 10 ml of 24 pentane. To this was added dropwise 5 ml of a pentane solution 25 containing bis(cyclopentadienyl) hafnium dimethyl (15 mg). After 26 stirring at room temperature for 15 minutes, the solvent was removed 27 in vacuo to leave a free-flowing supported catalyst.

Example 11

500 mg. of the supported catalyst of Example 10 was used to polymerize ethylene in a process similar to Example 1 except that the autoclave was pressured to 400 psig with ethylene and was heated to

- 1 80°C. The yield of linear granular, free-flowing polyethylene was 132 g.
- Although the invention has been described with reference to
- 4 its preferred embodiments, those of ordinary skill in the art may,
- 5 upon reading this disclosure, appreciate changes and modifications
- 6 which do not depart from the scope and spirit of the invention as
- 7 described above or claimed hereafter.

- Davison 952 silica (1 g) was slurried in pentane and treated with 1 ml of a 1 m solution of triethylaluminum
- 10 in hexane. The support was separated from the liquid by
- 11 decantation, washed twice with pentane and dried in vacuo.
- N,N-dimethylanilinium tetrakis
- .13 (pentafluorophenyl)boron (30 mg) was heated in toluene to
- 14 produce a homogeneous solution. The warm solution was added
- 15 dropwise to the support prepared as described above. The
- 16 solvent was removed in vacuo to leave a free-flowing
- 17 supported material.
- 18 This support material was slurried in 10 ml of
- 19 pentane. To this was added dropwise $10\ \mathrm{ml}$ of a pentane
- 20 solution containing 15 mg of rac-
- 21 dimethylsilylbis(tetrahydroindenyl) zirconium dimethyl.
- 22 After stirring at room temperature for 15 minutes, the
- 23 solvent was removed in vacuo to leave a free-flowing
- 24 supported catalyst.

-31-

Example 13

- 1 The catalyst of Example 12 (887 mg) was injected
- 2 by means of high-pressure nitrogen into a 1 liter stainless-
- 3 steel autoclave, which was previously flushed with nitrogen,
- 4 containing 300 ml dry, deoxygenated propylene at 40 degrees
- 5 C. After 20 minutes, the contents isolated the yield of
- 6 isotactic polypropylene was 101.7 g.

Example 14

- Davison 948 silica (3 g), dehydrated at 800
- degrees with an nitrogen purge was slurried in pentane and
- 9 treated with 8 ml of a 1.5 M solution of triethylaluminum in
- 10 hexane. The support was separated from the liquid by
- 11 decantation, washed six times with pentane, and dried in
- 12 vacuo.
- N, N-dimethylanilinium
- 14 tetrakis(pentafluorophenyl)baron (96 mg) was heated in 30 ml
- 15 warm toluene, to produce a homogeneous solution. The warm
- 16 solution was added dropwise to a toluene slurry of the
- 17 support prepared as described above. The solvent was
- 18 removed in vacuo with gentle heating to leave a free-flowing
- 19 supported material.
- This supported material was reslurried in pentane.
- 21 To this was added dropwise 20 ml of a pentane solution
- 22 containing 48 mg of bis(cyclopentadienyl)hafnium dimethyl.
- 23 After stirring at room temperature for 15 minutes, the
- 24 solvent was removed in vacuo to leave a free-flowing
- 25 supported catalyst.

The catalyst of Example 14 (100 mg) was injected with 100 ml hexane into a 1 liter stainless-steel autoclave, which was previously flushed with nitrogen, containing 300 ml of dry, deoxygenated hexane and pressurized with 200 psi ethylene at 60 degrees C. After 20 minutes the autoclave was cooled and vented and the contents isolated by filtration. The yield of linear polyethylene was 78.7 g.

ودبو

CLAIMS:

- A method for preparing a supported ionic catalyst comprising the steps of:
 - (a) combining, in a solvent or diluent
- (i) at least one metallocene component comprising a bis(cyclopentadienyl)metal compound containing at least one ligand capable of reacting with a proton, said metal being selected from the group consisting of titanium, zirconium and hafnium,
- (ii) at least one activator component comprising a cation capable of donating a proton and an anion, said anion being a single coordination complex comprising a plurality of lipophilic radicals covalently coordinated to and shielding a central charge-bearing metal or metalloid atom, said anion being bulky, labile and capable of stabilizing the metal cation formed as a result of reaction between the two, and

(iii) a catalyst support material suspended in a suitable solvent; and

- (b) recovering a supported catalyst product as a free-flowing solid or slurry.
- 2. The method of claim 1 wherein said activator component is represented by the following general formula:

$$[(L'-H)^{+}]_{d}[(M')^{m+}Q_{1}Q_{2}...Q_{n}]^{d-}$$

wherein:

L' is a neutral Lewis base; H is a hydrogen atom; [L'-H]⁺ is a Bronsted acid;

M' is a metal or metalloid selected from the groups subtended by Groups V-B to V-A of the Periodic Table of the Elements; i.e., Groups V-B, VI-B, VII-B, VIII, I-B, II-B, III-A, IV-A and V-A;

Q: to Qn are selected, independently, from the group consisting of hydride radicals, dialkylamido radicals, alkoxide and

aryloxide radicals, hydrocarbyl and substituted-hydrocarbyl radicals and organometalloid radicals and any one, but not more than one, of Q_1 to Q_n may be a halide radical with the remaining Q_1 to Q_n being, independently, selected from the foregoing radicals;

m is an integer from 1 to 7; n is an integer from 2 to 8; and n-m=d.

3. The method of claim 2 wherein said activator component is represented by the following general formula:

$$[L'-H]^+[BAr_1Ar_2X_3X_4]^-$$

wherein:

L' is a neutral Lewis base; H is a hydrogen atom; [L'-H]⁺ is a Bronsted acid; B is boron in a valence state of 3;

Ar and Ar are the same or different aromatic or substituted-aromatic hydrocarbon radicals, said radicals being optionally linked to each other through a stable bridging group; and X X_3 and X_4 are, independently, selected from the group consisting of hydride radicals, halide radicals, hydrocarbyl and substituted-hydrocarbyl radicals and organometalloid radicals.

- 4. The method of claim 3 wherein said bis(cyclopentadienyl)metal compound is represented by one of the following general formulae:
 - 1. $(A-Cp)HX_1X_2$
 - 2. (A-Cp)HX'1X'2
 - 3. (A-Cp)ML
 - 4. (Cp*)(CpR)HX,

Wherein:

M is a metal selected from the group consisting of titanium, zirconium and hafnium:

(A-Cp) comprises (Cp)(Cp*) or Cp-A'-Cp* and Cp and Cp* are the same or different substituted or unsubstituted cyclopentadienyl radicals;

A' is a covalent bridging group; L is an olefin, diolefin or aryne ligand;

X₁ and X₂ are, independently, selected from the group consisting of hydride radicals, hydrocarbyl radicals, substituted-hydrocarbyl radicals and organometalloid radicals;

X'₁ and X'₂ are joined and bound to the metal atom to form a metallacycle, in which the metal atom, X'₁ and X'₂ form a hydrocarbocyclic ring containing from about 3 to about 20 carbon atoms; and

R is a substituent on one of the cyclopentadienyl radicals which is also bound to the metal atom.

- 5. The method of claim 1 wherein said support is selected from alumina, silica, alumina-silica, talc, magnesia, zirconia, titania, finely divided polyolefins or mixtures thereof.
- The method of claim 5 wherein said support is alumina, silica or mixtures thereof.
- 7. The method of claim 4 wherein the metallocene is bis(cyclopentadienyl)zirconium dimethyl or bis(cyclopentadienyl)hafnium dimethyl; the activator is N,N-dimethylanilinium tetrakis(pentafluorophenyl)boron; and the support is alumina or silica.
- 8. The method in of claim 1, wherein the at least one metallocene and the at least one activator component are reacted in a solvent or diluent and the direct product or the decomposition product of said direct product is contacted in a suitable solvent with a

catalyst support material and thereafter the supported catalyst product is recovered as a free-flowing solid or slurry.

- 9. The method of claim 1, wherein the activator component is dissolved in an aromatic solvent and the support is slurried therein and allowed to react, the supported activator component is then contacted with the metallocene component in an aliphatic method.
- 10. A catalyst comprising the supported ionic catalyst obtained by the method of claim 1.

INTERNATIONAL SEARCH REPORT

International Application No PCT/US 90./07669.

I: CLASSI	IFICATION OF SUBJECT MATTER (it several classific	estion symbols apply, indicate all)	
	to International Patent Classification (IPC) or to both Natio	nal Classification and IPC	
IPC ⁵ :	C 08 F 4/76, 4/603, 10/00		
II. FIELDS	SEARCHED		· · · · · · · · · · · · · · · · · · ·
	Minimum Document		
Classificatio	on System C	lassification Symbols	
IPC ⁵		•	
IPC -	C 08 F	•	
	Documentation Searched other th	Alejmum Documentation	
	to the Extent that such Documents a	are included in the Fields Searched	
		•	
		<u> </u>	
III. DOCU	MENTS CONSIDERED TO BE RELEVANT	and the relevant passages 12	Relevant to Claim No. 13
ategory •	Citation of Document, 11 with Indication, where appr	opnate, of the relevant passages	
Α	EP, A, 0277004 (EXXON)		, 1
i	3 August 1988		
	see the claims		1
. !	cited in the application		: -
A	EP, A, 0232595 (EXXON)		1
_ ^	19 August 1987	•	
į.	see the claims		i
į	cited in the application	•	1
	0077000 (EVVON)		1 1
A	EP, A, 0277003 (EXXON) 3 August 1988		
	see the claims		
			
Α .	US, A, 4017525 (R.A. SEIT	ERQUIST)	. 1
	12 April 1977		
ļ	see the claims		
ļ			
ĺ			
1			·
A F i -	al categories of cited documents: 10	"T" later document published after	the International filing date
"A" doc	rument defining the general state of the art which is not	or priority date and not in conf cited to understand the princip	le or theory underlying the
con E" sari	sidered to be of particular relevance lier document but published on or after the international	invention "X" document of particular relevan	nce; the claimed invention
filin	ng date	cannot be considered novel o Involve an inventive step	
whi cita	ich is cited to establish the publication date of ending- tion or other special reason (as specified)	"Y" document of particular rejeval cannot be considered to involve	Y WE INVESTIGE SIGN ACCESS THE
"O" doc	cument referring to an oral disclosure, use, exhibition or ler means	document is combined with on ments, such combination being	
"P" dod	cument published prior to the international filing date but	in the art "&" document member of the same	patent family
	er than the priority date claimed		
	rIFICATION REPORT OF THE Actual Completion of the International Search	Date of Mailing of this International S	iearch Report
		1 3, 06, 91	. /
	th April 1991	Signature of Authorized Officer	A. Lactor
Internation	nal Searching Authority	/	May Ill He
•	EUROPEAN PATENT OFFICE	miss T. MORTENSEN/	

ANNEX TO THE INTERNATIONAL SEARCH REPORT ON INTERNATIONAL PATENT APPLICATION NO.

US 9007669

44096 SA

This annex lists the patent family members relating to the patent documents cited in the above-mentioned international search report. The members are as contained in the European Patent Office EDP file on 04/06/91

The European Patent Office is in no way liable for these particulars which are merely given for the purpose of information.

Patent document cited in search report	Publication date	Patent family member(s)		Publication date
EP-A- 0277004	03-08-88	AU-A- JP-T- WO-A-	1245288 1502036 8805793	24-08-88 13-07-89 11-08-88
EP-A- 0232595	19-08-87	US-A- AU-B- AU-A- EP-A- JP-T- WO-A-	4701432 599622 6728587 0245482 63501369 8702991	20-10-87 26-07-90 02-06-87 19-11-87 26-05-88 21-05-87
EP-A- 0277003	03-08-88	AU-A- JP-T- WO-A-	1294588 1501950 8805792	24-08-88 .06-07-89 11-08-88
US-A- 4017525	12-04-77	None		